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I, KIM MARSHALL, MANAGER EXAMINATION SUPPORT AND SALES,
hereby certify that the annexed is a true copy of the Provisional specification in
connection with Application No. PO 6972 for a patent by THE COUNCIL OF THE
QUEENSLAND INSTITUTE OF MEDICAL RESEARCH filed on 23 May 1997

I further certify that the annexed specification is not, as yet, open to public inspection.

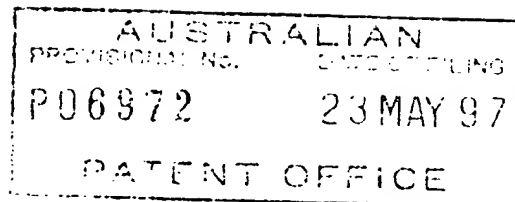
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WITNESS my hand this First
day of June 1998

KIM MARSHALL
MANAGER EXAMINATION SUPPORT AND
SALES



Regulation 3.2



The Council of The Queensland Institute of Medical Research

A U S T R A L I A
Patents Act 1990

PROVISIONAL SPECIFICATION

for the invention entitled:

"A novel gene and uses therefor"

The invention is described in the following statement:

- 1A -

A NOVEL GENE AND USES THEREFOR

5 The present invention relates generally to a novel human gene and to derivatives and mammalian, animal, insect, nematodes, avian and microbial homologues thereof. The present invention further provides pharmaceutical compositions and diagnostic agents as well as genetic molecules useful in gene replacement therapy and recombinant molecules useful in protein replacement therapy.

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Throughout this specification and the claims which follow, unless the context requires otherwise, the word "comprise", or variations such as "comprises" or "comprising", will be understood to imply the inclusion of a stated integer or group of integers but not the exclusion of any other integer or group of integers.

15

Sequence Identity Numbers (SEQ ID NOS.) for the nucleotide and amino acid sequences referred to in the specification are defined at the end of the description.

The increasing sophistication of recombinant DNA technology is greatly facilitating research
20 and development in the medical and allied health fields. There is growing need to develop recombinant and genetic molecules for use in diagnosis, conventional pharmaceutical preparations as well as gene and protein replacement therapies.

In work leading up to the present invention, the inventors sought to identify and clone human
25 genes which might be useful as potential diagnostic and/or therapeutic agents. One area of particular interest is in the field of gene regulators.

Gene expression generally requires interaction between a regulatory protein and an appropriate recognition sequence of a target gene. Regulatory proteins comprise in many
30 cases a domain or motif that facilitates binding to DNA. One particular motif comprises

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small sequence units repeated in tandem with each unit folded about a zinc atom to form separate structural domains. This motif is now referred to as a zinc finger domain. Such a domain is generally defined by the number of cysteine (C) and histidine (H) residues.

- 5 In accordance with the present invention, a gene has been identified from the human genome with an N-terminal region resembling a zinc-finger domain of a novel type.

Accordingly, one aspect of the present invention contemplates an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence 10 encoding an amino acid sequence having homology to a regulator of gene expression or a derivative of said gene regulator.

More particularly the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides encoding or complementary to a sequence encoding putative 15 regulator of gene expression wherein said regulator comprises a zinc finger domain of an $(HC_3)_2$ type.

Even more particularly, the present invention is directed to an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

20

- (i) a nucleotide sequence set forth in SEQ ID NO:1;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:2;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- 25 (iv) a nucleotide sequence capable of hybridizing under low stringency conditions to the nucleotide sequence set forth in (i), (ii) or (iii).

In a related embodiment, the present invention provides an isolated nucleic acid molecule comprising a sequence of nucleotides or a complementary form thereof selected from:

30

- (i) a nucleotide sequence set forth in SEQ ID NO:3;
- (ii) a nucleotide sequence encoding an amino acid sequence set forth in SEQ ID NO:4;
- (iii) a nucleotide sequence having at least about 40% similarity to the nucleotide sequence of (i) or (ii); and
- 5 (iv) a nucleotide sequence capable of hybridizing under low stringency conditions to the nucleotide sequence set forth in (i), (ii) or (iii).

Preferably, the percentage similarity is at least about 50%. More preferably, the percentage similarity is at least about 60%.

10

Reference herein to a low stringency at 42°C includes and encompasses from at least about 1% v/v to at least about 15% v/v formamide and from at least about 1M to at least about 2M salt for hybridisation, and at least about 1M to at least about 2M salt for washing conditions. Alternative stringency conditions may be applied where necessary, such as medium stringency, which 15 includes and encompasses from at least about 16% v/v to at least about 30% v/v formamide and from at least about 0.5M to at least about 0.9M salt for hybridisation, and at least about 0.5M to at least about 0.9M salt for washing conditions, or high stringency, which includes and encompasses from at least about 31% v/v to at least about 50% v/v formamide and from at least about 0.01M to at least about 0.15M salt for hybridisation, and at least about 0.01M to at least 20 about 0.15M salt for washing conditions.

The term "similarity" as used herein includes exact identity between compared sequences at the nucleotide or amino acid level. Where there is non-identity at the nucleotide level, "similarity" includes differences between sequences which result in different amino acids that are nevertheless 25 related to each other at the structural, functional, biochemical and/or conformational levels. Where there is non-identity at the amino acid level, "similarity" includes amino acids that are nevertheless related to each other at the structural, functional, biochemical and/or conformational levels.

30 The present invention extends to nucleic acid molecules with percentage similarities of approximately 65%, 70%, 75%, 80%, 85%, 90% or 95% or above or a percentage in between.

The nucleic acid molecule of the present invention is hereinafter referred to as constituting the "mcg4" gene. The protein encoded by *mcg4* is referred to herein as "MCG4". The *mcg4* gene is proposed to encode, in accordance with the present invention, a regulator of gene expression and to comprise the novel zinc finger domain (HC₃)₂. A regulator of gene 5 expression includes a transcription factor. Regulation may be at the level of nucleic acid:protein or protein:protein interaction.

The present invention extends to the naturally occurring genomic *mcg4* nucleotide sequence or corresponding cDNA sequence or to derivatives thereof. Derivatives contemplated in the 10 present invention include fragments, parts, portions, mutants, homologues and analogues of MCG4 or the corresponding genetic sequence. Derivatives also include single or multiple amino acid substitutions, deletions and/or additions to MCG4 or single or multiple nucleotide substitutions, deletions and/or additions to *mcg4*. "Additions" to the amino acid or nucleotide sequences include fusions with other peptides, polypeptides or proteins or fusions to 15 nucleotide sequences. Reference herein to "MCG4" or "*mcg4*" includes references to all derivatives thereof including functional derivatives and immunologically interactive derivatives of MCG4.

The *mcg4* of the present invention is particularly exemplified herein from humans and in 20 particular from human chromosome 11q13.

The present invention extends, however, to a range of homologues from, for example, primates, livestock animals (eg. sheep, cows, horses, donkeys, pigs), companion animals (eg. dogs, cats) laboratory test animals (eg. rabbits, mice, rats, guinea pigs), birds (eg. chickens, 25 ducks, geese, parrots), insects, nematodes, eukaryotic microorganisms and captive wild animals (eg. deer, foxes, kangaroos). Reference herein to *mcg4* or MCG4 includes reference to these molecules of human origin as well as novel forms of non-human origin.

The nucleic acid molecules of the present invention may be DNA or RNA. When the nucleic 30 acid molecule is in DNA form, it may be genomic DNA or cDNA. RNA forms of the nucleic

acid molecules of the present invention are generally mRNA.

Although the nucleic acid molecules of the present invention are generally in isolated form, they may be integrated into or ligated to or otherwise fused or associated with other genetic 5 molecules such as vector molecules and in particular expression vector molecules. Vectors and expression vectors are generally capable of replication and, if applicable, expression in one or both of a prokaryotic cell or a eukaryotic cell. Preferably, prokaryotic cells include *E. coli*, *Bacillus sp* and *Pseudomonas sp*. Preferred eukaryotic cells include yeast, fungal, mammalian and insect cells.

10

Accordingly, another aspect of the present invention contemplates a genetic construct comprising a vector portion and an animal, more particularly a mammalian and even more particularly a human *mcg4* gene portion, which *mcg4* gene portion is capable of encoding an MCG4 polypeptide or a functional or immunologically interactive derivative thereof.

15

Preferably, the *mcg4* gene portion of the genetic construct is operably linked to a promoter on the vector such that said promoter is capable of directing expression of said *mcg4* gene portion in an appropriate cell.

20 In addition, the *mcg4* gene portion of the genetic construct may comprise all or part of the gene fused to another genetic sequence such as a nucleotide sequence encoding glutathione-S-transferase or part thereof.

The present invention extends to such genetic constructs and to prokaryotic or eukaryotic cells
25 comprising same.

It is proposed in accordance with the present invention that MCG4 is a transcription factor involved in gene regulation. Mutations in *mcg4* may result in aberrations in gene regulation leading to the development of or a propensity to develop various types of cancer. In this
30 regard, although not wishing to limit the present invention to any one hypothesis or mode of action, it is proposed that *mcg4* or its expression product may be involved in the tissue-

specific or temporal regulation of particular genes.

A deletion or aberration in the *mcg4* gene may also be important in the detection of cancer or a propensity to develop cancer. An aberration may be a homozygous mutation or a 5 heterozygous mutation. The detection may occur at the foetal or post-natal level. Detection may also be at the germline or somatic cell level. Furthermore, a risk of developing cancer may be determined by assaying for aberrations in the parents and/or proband of a subject under investigation.

10 According to this aspect of the present invention, there is contemplated a method of detecting a condition caused or facilitated by an aberration in *mcg4*, said method comprising determining the presence of a single or multiple nucleotide substitution, deletion and/or addition or other aberration to one or both alleles of said *mcg4* wherein the presence of such a nucleotide substitution, deletion and/or addition or other aberration may be indicative of 15 said condition or a propensity to develop said condition.

The nucleotide substitutions, additions or deletions may be detected by any convenient means including nucleotide sequencing, restriction fragment length polymorphism (RFLP), polymerase chain reaction (PCR), oligonucleotide hybridization and single stranded 20 conformation polymorphism analysis (SSCP) amongst many others. An aberration includes modification to existing nucleotides such as to modify glycosylation signal amongst other effects.

In an alternative method, aberrations in the *mcg4* gene are detected by screening for mutations 25 in MCG4.

A mutation in MCG4 may be a single or multiple amino acid substitution, addition and/or deletion. The mutation in *mcg4* may also result in either no translation product being produced or a product in truncated form. A mutant may also be an altered glycosylation 30 pattern or the introduction of side chain modifications to amino acid residues.

According to this aspect of the present invention, there is provided a method of detecting a condition caused or facilitated by an aberration in *mcg4*, said method comprising screening for a single or multiple amino acid substitution, deletion and/or addition to MCG4 wherein the presence of such a mutation is indicative of or a propensity to develop said condition.

5

A particularly convenient means of detecting a mutation in MCG4 is by use of antibodies.

Accordingly another aspect of the present invention is directed to antibodies to MCG4 and its derivatives. Such antibodies may be monoclonal or polyclonal and may be selected from 10 naturally occurring antibodies to MCG4 or may be specifically raised to MCG4 or derivatives thereof. In the case of the latter, MCG4 or its derivatives may first need to be associated with a carrier molecule. The antibodies to MCG4 of the present invention are particularly useful as diagnostic agents.

15 For example, antibodies to MCG4 and its derivatives can be used to screen for wild-type MCG4 or for mutated MCG4 molecules. The latter may occur, for example, during or prior to certain cancer development. A differential binding assay is also particularly useful. Techniques for such assays are well known in the art and include, for example, sandwich assays and ELISA. Knowledge of normal MCG4 levels or the presence of wild-type MCG4 may be important for 20 diagnosis of certain cancers or a predisposition for development of cancers or for monitoring certain therapeutic protocols.

As stated above antibodies to MCG4 of the present invention may be monoclonal or polyclonal or may be fragments of antibodies such as Fab fragments. Furthermore, the present invention 25 extends to recombinant and synthetic antibodies and to antibody hybrids. A "synthetic antibody" is considered herein to include fragments and hybrids of antibodies.

For example, specific antibodies can be used to screen for wild-type MCG4 molecule or specific mutant molecules such as molecules having a certain deletion. This would be important, for 30 example, as a means for screening for levels of MCG4 in a cell extract or other biological fluid or purifying MCG4 made by recombinant means from culture supernatant fluid or purified from

a cell extract. Techniques for the assays contemplated herein are known in the art and include, for example, sandwich assays and ELISA.

It is within the scope of this invention to include any second antibodies (monoclonal, polyclonal 5 or fragments of antibodies or synthetic antibodies) directed to the first mentioned antibodies discussed above. Both the first and second antibodies may be used in detection assays or a first antibody may be used with a commercially available anti-immunoglobulin antibody. An antibody as contemplated herein includes any antibody specific to any region of wild-type MCG4 or to a specific mutant phenotype or to a deleted or otherwise altered region.

10

Both polyclonal and monoclonal antibodies are obtainable by immunization of a suitable animal or bird with MCG4 or its derivatives and either type is utilizable for immunoassays. The methods of obtaining both types of sera are well known in the art. Polyclonal sera are less preferred but are relatively easily prepared by injection of a suitable laboratory animal or bird 15 with an effective amount of MCG4 or antigenic parts thereof or derivatives thereof, collecting serum from the animal or bird, and isolating specific sera by any of the known immunoabsorbent techniques. Although antibodies produced by this method are utilizable in virtually any type of immunoassay, they are generally less favoured because of the potential heterogeneity of the product.

20

The use of monoclonal antibodies in an immunoassay is particularly preferred because of the ability to produce them in large quantities and the homogeneity of the product. The preparation of hybridoma cell lines for monoclonal antibody production derived by fusing an immortal cell line and lymphocytes sensitized against the immunogenic preparation can be done by techniques 25 which are well known to those who are skilled in the art.

Another aspect of the present invention contemplates a method for detecting MCG4 or a derivative thereof in a biological sample said method comprising contacting said biological sample with an antibody specific for MCG4 or its derivatives or homologues for a time and under 30 conditions sufficient for an antibody-MCG4 complex to form, and then detecting said complex.

Preferably, the biological sample is a cell extract from a human or other animal or a bird.

The presence of MCG4 may be accomplished in a number of ways such as by Western blotting and ELISA procedures. A wide range of immunoassay techniques are available as can be seen
5 by reference to US Patent Nos. 4,016,043, 4, 424,279 and 4,018,653. These include both single-site and two-site or "sandwich" assays of the non-competitive types, as well as traditional competitive binding assays. These assays also include direct binding of a labelled antibody to a target.

- 10 Sandwich assays are among the most useful and commonly used assays and are favoured for use in the present invention. A number of variations of the sandwich assay technique exist, and all are intended to be encompassed by the present invention. Briefly, in a typical forward assay, an unlabelled antibody is immobilized on a solid substrate and the sample to be tested brought into contact with the bound molecule. After a suitable period of incubation, for a period of time
15 sufficient to allow formation of an antibody-antigen complex, a second antibody specific to the antigen, labelled with a reporter molecule capable of producing a detectable signal is then added and incubated, allowing time sufficient for the formation of another complex of antibody-antigen-labelled antibody. Any unreacted material is washed away, and the presence of the antigen is determined by observation of a signal produced by the reporter molecule. The results may either
20 be qualitative, by simple observation of the visible signal, or may be quantitated by comparing with a control sample containing known amounts of hapten. Variations on the forward assay include a simultaneous assay, in which both sample and labelled antibody are added simultaneously to the bound antibody. These techniques are well known to those skilled in the art, including any minor variations as will be readily apparent. In accordance with the present
25 invention the sample is one which might contain MCG4 including cell extract or, tissue biopsy. The sample is, therefore, generally a biological sample comprising biological fluid but also extends to fermentation fluid and supernatant fluid such as from a cell culture.

- In the typical forward sandwich assay, a first antibody having specificity for the MCG4 or an
30 antigenic part thereof or a derivative thereof or antigenic parts thereof, is either covalently or passively bound to a solid surface. The solid surface is typically glass or a polymer, the most

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commonly used polymers being cellulose, polyacrylamide, nylon, polystyrene, polyvinyl chloride or polypropylene. The solid supports may be in the form of tubes, beads, discs of microplates, or any other surface suitable for conducting an immunoassay. The binding processes are well-known in the art and generally consist of cross-linking covalently binding or physically adsorbing, 5 the polymer-antibody complex is washed in preparation for the test sample. An aliquot of the sample to be tested is then added to the solid phase complex and incubated for a period of time sufficient (e.g. 2-40 minutes) and under suitable conditions (e.g. 25 °C) to allow binding of any subunit present in the antibody. Following the incubation period, the antibody subunit solid phase is washed and dried and incubated with a second antibody specific for a portion of the 10 hapten. The second antibody is linked to a reporter molecule which is used to indicate the binding of the second antibody to the hapten.

An alternative method involves immobilizing the target molecules in the biological sample and then exposing the immobilized target to specific antibody which may or may not be labelled with 15 a reporter molecule. Depending on the amount of target and the strength of the reporter molecule signal, a bound target may be detectable by direct labelling with the antibody. Alternatively, a second labelled antibody, specific to the first antibody is exposed to the target-first antibody complex to form a target-first antibody-second antibody tertiary complex. The complex is detected by the signal emitted by the reporter molecule.

20

By "reporter molecule" as used in the present specification, is meant a molecule which, by its chemical nature, provides an analytically identifiable signal which allows the detection of antigen-bound antibody. Detection may be either qualitative or quantitative. The most commonly used reporter molecules in this type of assay are either enzymes, fluorophores or radionuclide 25 containing molecules (i.e. radioisotopes) and chemiluminescent molecules.

In the case of an enzyme immunoassay, an enzyme is conjugated to the second antibody, generally by means of glutaraldehyde or periodate. As will be readily recognized, however, a wide variety of different conjugation techniques exist, which are readily available to the skilled artisan. Commonly used enzymes include horseradish peroxidase, glucose oxidase, beta-30 galactosidase and alkaline phosphatase, amongst others. The substrates to be used with the specific enzymes are generally chosen for the production, upon hydrolysis by the corresponding

enzyme, of a detectable colour change. Examples of suitable enzymes include alkaline phosphatase and peroxidase. It is also possible to employ fluorogenic substrates, which yield a fluorescent product rather than the chromogenic substrates noted above. In all cases, the enzyme-labelled antibody is added to the first antibody hapten complex, allowed to bind, and 5 then the excess reagent is washed away. A solution containing the appropriate substrate is then added to the complex of antibody-antigen-antibody. The substrate will react with the enzyme linked to the second antibody, giving a qualitative visual signal, which may be further quantitated, usually spectrophotometrically, to give an indication of the amount of hapten which was present in the sample. "Reporter molecule" also extends to use of cell agglutination or inhibition of 10 agglutination such as red blood cells on latex beads, and the like.

Alternately, fluorescent compounds, such as fluorescein and rhodamine, may be chemically coupled to antibodies without altering their binding capacity. When activated by illumination with light of a particular wavelength, the fluorochrome-labelled antibody adsorbs the light 15 energy, inducing a state to excitability in the molecule, followed by emission of the light at a characteristic colour visually detectable with a light microscope. As in the EIA, the fluorescent labelled antibody is allowed to bind to the first antibody-hapten complex. After washing off the unbound reagent, the remaining tertiary complex is then exposed to the light of the appropriate wavelength the fluorescence observed indicates the presence of the hapten of interest. 20 Immunofluorescence and EIA techniques are both very well established in the art and are particularly preferred for the present method. However, other reporter molecules, such as radioisotope, chemiluminescent or bioluminescent molecules, may also be employed.

As stated above, the present invention extends to genetic constructs capable of encoding 25 MCG4 or functional derivatives thereof. Such genetic constructs are also contemplated to be useful in modulating expression of specific genes in which *mcg4* is involved in tissue-specific or temporal regulation.

Accordingly, another aspect of the present invention is directed to a genetic construct 30 comprising a nucleotide sequence encoding a peptide, polypeptide or protein and *mcg4* or a functional derivative or homologue thereof capable of modulating the expression of said

nucleotide sequence.

The present invention is further described with reference to the following non-limiting Figures and Examples.

5

In the Figures:

Figure 1 is a representation of the nucleotide sequence and corresponding amino acid sequence of *mcg4*.

10

Figure 2 is a representation of the alignment of the human MCG4 amino acid sequence with a translation of a partial murine expressed sequence tag (EST).

15 Figure 3 is a representation of the alignment of the human MCG4 amino acid sequence with a translation of a partial nematode EST.

Figure 4 is a diagrammatic representation showing a predicted structure of MCG4.

Figure 5 is a representation of sensitive sequence homology search of related cysteine-20 containing motifs in another *Caenorhabditis elegans* protein.

Figure 6 is a representation showing that a related cysteine containing motif is present in the GATA-binding transcription factor from *Saccharomyces pombe*.

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EXAMPLE 1

A human gene (designated *mcg4*) was identified on chromosome 11q13 that on the basis of sequence homology is predicted to encode a putative transcription factor of 310 amino acids
5 (Fig. 1). *mcg4* is transcribed as an ~ 1.6kb mRNA.

EXAMPLE 2

The expressed sequence tag (EST) database contains partial sequence data for the murine (Fig.
10 2) and nematode (Fig. 3) homologues.

EXAMPLE 3

MCG4 contains a sequence of cysteine residues within the N-terminal region of the protein
15 that resembles zinc-finger binding domains of a novel type, ie. $(HC_3)_2$ [Fig. 4].

EXAMPLE 4

Sensitive sequence homology searches reveal that related cysteine-containing motifs are
20 present in another *C. elegans* protein (Fig. 5) as well as the GATA-binding transcription
factor from *S. pombe* (Fig. 6).

EXAMPLE 5

25 *mcg4* will have commercial value due to its likelihood of encoding a novel transcription factor
that is highly conserved amongst organisms, thus suggesting an integral role in gene
regulation. *mcg4* may also be involved in some way in tissue-specific or temporal regulation
of certain genes, thus making it a potential target for modulating expression of those
downstream effectors.

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Those skilled in the art will appreciate that the invention described herein is susceptible to variations and modifications other than those specifically described. It is to be understood that the invention includes all such variations and modifications. The invention also includes all of the steps, features, compositions and compounds referred to or indicated in this specification,
5 individually or collectively, and any and all combinations of any two or more of said steps or features.

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SEQUENCE LISTING

(1) GENERAL INFORMATION:

(i) APPLICANT: The Council of The Queensland Institute of Medical Research

(ii) TITLE OF INVENTION: A NOVEL GENE AND USES THEREFOR

(iii) NUMBER OF SEQUENCES: 2

(iv) CORRESPONDENCE ADDRESS:

- (A) ADDRESSEE: DAVIES COLLISON CAVE
- (B) STREET: 1 LITTLE COLLINS STREET
- (C) CITY: MELBOURNE
- (D) STATE: VICTORIA
- (E) COUNTRY: AUSTRALIA
- (F) ZIP: 3000

(v) COMPUTER READABLE FORM:

- (A) MEDIUM TYPE: Floppy disk
- (B) COMPUTER: IBM PC compatible
- (C) OPERATING SYSTEM: PC-DOS/MS-DOS
- (D) SOFTWARE: PatentIn Release #1.0, Version #1.25

(vi) CURRENT APPLICATION DATA:

- (A) APPLICATION NUMBER: AUSTRALIAN PROVISIONAL
- (B) FILING DATE:
- (C) CLASSIFICATION:

(viii) ATTORNEY/AGENT INFORMATION:

- (A) NAME: HUGHES, DR E JOHN L
- (C) REFERENCE/DOCKET NUMBER: EJH/AF

(ix) TELECOMMUNICATION INFORMATION:

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- (B) TELEFAX: +61 3 9254 2770
- (C) TELEX: AA 31787

(2) INFORMATION FOR SEQ ID NO:1:

- (i) SEQUENCE CHARACTERISTICS:

 - (A) LENGTH: 1242 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: DNA

- (ix) FEATURE:
 (A) NAME/KEY: CDS
 (B) LOCATION: 30..959

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

TCAGTAAACA CAGAGACTGG GGATCGATC ATG GGG CTT TGT AAG TGC CCC AAG Met Gly Leu Cys Lys Cys Pro Lys	53
1 5	
AGA AAG GTG ACC AAC CTG TTC TGC TTC GAA CAT CGG GTC AAC GTC TGC Arg Lys Val Thr Asn Leu Phe Cys Phe Glu His Arg Val Asn Val Cys	101
10 15 20	
GAG CAC TGC CTG GTA GCC AAT CAC GCC AAG TGC ATC GTC CAG TCC TAC Glu His Cys Leu Val Ala Asn His Ala Lys Cys Ile Val Gln Ser Tyr	149
25 30 35 40	
CTG CAA TGG CTC CAA GAT AGC GAC TAC AAC CCC AAT TGC CGC CTG TGC Leu Gln Trp Leu Gln Asp Ser Asp Tyr Asn Pro Asn Cys Arg Leu Cys	197
45 50 55	
AAC ATA CCC CTG GCC AGC CGA GAG ACG ACC CGC CTT GTC TGC TAT GAT Asn Ile Pro Leu Ala Ser Arg Glu Thr Thr Arg Leu Val Cys Tyr Asp	245
60 65 70	
CTC TTT CAC TGG GCC TGC CTC AAT GAA CGT GCT GCC CAG CTA CCC CGA Leu Phe His Trp Ala Cys Leu Asn Glu Arg Ala Ala Gln Leu Pro Arg	293
75 80 85	
AAC ACG GCA CCT GCC GGC TAT CAG TGC CCC AGC TGC AAT GGC CCC ATC Asn Thr Ala Pro Ala Gly Tyr Gln Cys Pro Ser Cys Asn Gly Pro Ile	341
90 95 100	
TTC CCC CCA ACC AAC CTG GCT GGC CCC GTG GCC TCC GCA CTG AGA GAG Phe Pro Pro Thr Asn Leu Ala Gly Pro Val Ala Ser Ala Leu Arg Glu	389
105 110 115 120	
AAG CTG GCC ACA GTC AAC TGG GCC CGG GCA GGA CTG GGC CTC CCT CTG Lys Leu Ala Thr Val Asn Trp Ala Arg Ala Gly Leu Gly Leu Pro Leu	437
125 130 135	
ATC GAT GAG GTG GTG AGC CCA GAG CCC GAG CCC CTC AAC ACG TCT GAC Ile Asp Glu Val Val Ser Pro Glu Pro Glu Pro Leu Asn Thr Ser Asp	485
140 145 150	
TTC TCT GAC TGG TCT AGT TTT AAT GCC AGC AGT ACC CCT GGA CCA GAG Phe Ser Asp Trp Ser Ser Phe Asn Ala Ser Ser Thr Pro Gly Pro Glu	533
155 160 165	
GAG GTA GAC AGC GCC TCT GCT GCC CCA GCC TTC TAC AGC CAG GCC CCC Glu Val Asp Ser Ala Ser Ala Ala Pro Ala Phe Tyr Ser Gln Ala Pro	581
170 175 180	

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CGG CCC CCA GCT TCC CCA GGC CGG CCC GAG CAG CAC ACA GTG ATC CAC Arg Pro Pro Ala Ser Pro Gly Arg Pro Glu Gln His Thr Val Ile His 185 190 195 200	629
ATG GGC AAT CCT GAG CCC TTG ACT CAC GCC CCT AGG AAG GTG TAT GAT Met Gly Asn Pro Glu Pro Leu Thr His Ala Pro Arg Lys Val Tyr Asp 205 210 215	677
ACG CGG GAT GAT GAC CGG ACA CCA GGC CTC CAT GGA GAC TGT GAC GAT Thr Arg Asp Asp Asp Arg Thr Pro Gly Leu His Gly Asp Cys Asp Asp 220 225 230	725
GAC AAG TAC CGA CGT CGG CCG GCC TTG GGT TGG CTG GCC CGG CTG CTA Asp Lys Tyr Arg Arg Pro Ala Leu Gly Trp Leu Ala Arg Leu Leu 235 240 245	773
AGG AGC CGG GCT GGG TCT CGG AAG CGA CCG CTG ACC CTG CTC CAG CGG Arg Ser Arg Ala Gly Ser Arg Lys Arg Pro Leu Thr Leu Leu Gln Arg 250 255 260	821
GCG GGG CTG CTG CTA CTC TTG GGA CTG CTG GGC TTC CTG GCC CTC CTT Ala Gly Leu Leu Leu Leu Gly Leu Leu Gly Phe Leu Ala Leu Leu 265 270 275 280	869
GCC CTC ATG TCT CGC CTA GGC CGG GCC GCA GCT GAC AGC GAT CCC AAC Ala Leu Met Ser Arg Leu Gly Arg Ala Ala Ala Asp Ser Asp Pro Asn 285 290 295	917
CTG GAC CCA CTC ATG AAC CCT CAC ATC CGC GTG GGC CCC TCC TGA Leu Asp Pro Leu Met Asn Pro His Ile Arg Val Gly Pro Ser * 300 305 310	962
GCCCCCTTGC TTGTGGCTAG GCCAGCCTAG GATGTGGTT CTGTGGAGGA GAGGCAGGGT AATGGGGAGG CTGAGGGCAC CTCTTCACTG CCCCTCTCCC TCAAGCCTAA GACACTAAGA CCCCAGACCC AAAGCCAAGT CCACCAGAGT GGCTCGCAGG CCAGGCCTGG AGTCCCCGTG GGTCAAGCAT TTGTCTTGAC TTGCTTCTC CCGGGTCTCC AGCCTCCGAC CCCTCGCCCC ATGAAGGAGC TGGCAGGTGG AAATAAACAA CAACTTTATT	1022 1082 1142 1202 1242

(2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 310 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Gly Leu Cys Lys Cys Pro Lys Arg Lys Val Thr Asn Leu Phe Cys 1 5 10 15
Phe Glu His Arg Val Asn Val Cys Glu His Cys Leu Val Ala Asn His 20 25 30
Ala Lys Cys Ile Val Gln Ser Tyr Leu Gln Trp Leu Gln Asp Ser Asp 35 40 45
Tyr Asn Pro Asn Cys Arg Leu Cys Asn Ile Pro Leu Ala Ser Arg Glu 50 55 60

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Thr Thr Arg Leu Val Cys Tyr Asp Leu Phe His Trp Ala Cys Leu Asn
65 70 75 80

Glu Arg Ala Ala Gln Leu Pro Arg Asn Thr Ala Pro Ala Gly Tyr Gln
85 90 95

Cys Pro Ser Cys Asn Gly Pro Ile Phe Pro Pro Thr Asn Leu Ala Gly
100 105 110

Pro Val Ala Ser Ala Leu Arg Glu Lys Leu Ala Thr Val Asn Trp Ala
115 120 125

Arg Ala Gly Leu Gly Leu Pro Leu Ile Asp Glu Val Val Ser Pro Glu
130 135 140

Pro Glu Pro Leu Asn Thr Ser Asp Phe Ser Asp Trp Ser Ser Phe Asn
145 150 155 160

Ala Ser Ser Thr Pro Gly Pro Glu Glu Val Asp Ser Ala Ser Ala Ala
165 170 175

Pro Ala Phe Tyr Ser Gln Ala Pro Arg Pro Pro Ala Ser Pro Gly Arg
180 185 190

Pro Glu Gln His Thr Val Ile His Met Gly Asn Pro Glu Pro Leu Thr
195 200 205

His Ala Pro Arg Lys Val Tyr Asp Thr Arg Asp Asp Arg Thr Pro
210 215 220

Gly Leu His Gly Asp Cys Asp Asp Asp Lys Tyr Arg Arg Arg Pro Ala
225 230 235 240

Leu Gly Trp Leu Ala Arg Leu Leu Arg Ser Arg Ala Gly Ser Arg Lys
245 250 255

Arg Pro Leu Thr Leu Leu Gln Arg Ala Gly Leu Leu Leu Leu Gly
260 265 270

Leu Leu Gly Phe Leu Ala Leu Leu Ala Leu Met Ser Arg Leu Gly Arg
275 280 285

Ala Ala Ala Asp Ser Asp Pro Asn Leu Asp Pro Leu Met Asn Pro His
290 295 300

Ile Arg Val Gly Pro Ser
305 310

DATED this 23rd day of May 1997

The Council of The Queensland Institute of Medical Research
By DAVIES COLLISON CAVE
Patent Attorneys for the Applicants

FIGURE 1

TCAGTAAACA CAGAGACTGG GGATCGATC ATG GGG CTT TGT AAG TGC CCC AAG Met Gly Leu Cys Lys Cys Pro Lys	53
1 5	
AGA AAG GTG ACC AAC CTG TTC TGC TTC GAA CAT CGG GTC AAC GTC TGC Arg Lys Val Thr Asn Leu Phe Cys Phe Glu His Arg Val Asn Val Cys	101
10 15 20	
GAG CAC TGC CTG GTA GCC AAT CAC GCC AAG TGC ATC GTC CAG TCC TAC Glu His Cys Leu Val Ala Asn His Ala Lys Cys Ile Val Gln Ser Tyr	149
25 30 35 40	
CTG CAA TGG CTC CAA GAT AGC GAC TAC AAC CCC AAT TGC CGC CTG TGC Leu Gln Trp Leu Gln Asp Ser Asp Tyr Asn Pro Asn Cys Arg Leu Cys	197
45 50 55	
AAC ATA CCC CTG GCC AGC CGA GAG ACG ACC CGC CTT GTC TGC TAT GAT Asn Ile Pro Leu Ala Ser Arg Glu Thr Thr Arg Leu Val Cys Tyr Asp	245
60 65 70	
CTC TTT CAC TGG GCC TGC CTC AAT GAA CGT GCT GCC CAG CTA CCC CGA Leu Phe His Trp Ala Cys Leu Asn Glu Arg Ala Ala Gln Leu Pro Arg	293
75 80 85	
AAC ACG GCA CCT GCC GGC TAT CAG TGC CCC AGC TGC AAT GGC CCC ATC Asn Thr Ala Pro Ala Gly Tyr Gln Cys Pro Ser Cys Asn Gly Pro Ile	341
90 95 100	
TTC CCC CCA ACC AAC CTG GCT GGC CCC GTG GCC TCC GCA CTG AGA GAG Phe Pro Pro Thr Asn Leu Ala Gly Pro Val Ala Ser Ala Leu Arg Glu	389
105 110 115 120	
AAG CTG GCC ACA GTC AAC TGG GCC CGG GCA GGA CTG GGC CTC CCT CTG Lys Leu Ala Thr Val Asn Trp Ala Arg Ala Gly Leu Gly Leu Pro Leu	437
125 130 135	
ATC GAT GAG GTG GTG AGC CCA GAG CCC GAG CCC CTC AAC ACG TCT GAC Ile Asp Glu Val Val Ser Pro Glu Pro Glu Pro Leu Asn Thr Ser Asp	485
140 145 150	
TTC TCT GAC TGG TCT AGT TTT AAT GCC AGC AGT ACC CCT GGA CCA GAG Phe Ser Asp Trp Ser Ser Phe Asn Ala Ser Ser Thr Pro Gly Pro Glu	533
155 160 165	
GAG GTA GAC AGC GCC TCT GCT GCC CCA GCC TTC TAC AGC CAG GCC CCC Glu Val Asp Ser Ala Ser Ala Ala Pro Ala Phe Tyr Ser Gln Ala Pro	581
170 175 180	
CGG CCC CCA GCT TCC CCA GGC CGG CCC GAG CAG CAC ACA GTG ATC CAC Arg Pro Pro Ala Ser Pro Gly Arg Pro Glu Gln His Thr Val Ile His	629
185 190 195 200	
ATG GGC AAT CCT GAG CCC TTG ACT CAC GCC CCT AGG AAG GTG TAT GAT Met Gly Asn Pro Glu Pro Leu Thr His Ala Pro Arg Lys Val Tyr Asp	677
205 210 215	

Figure 1 (continued)

ACG CGG GAT GAT GAC CGG ACA CCA GGC CTC CAT GGA GAC TGT GAC GAT Thr Arg Asp Asp Asp Arg Thr Pro Gly Leu His Gly Asp Cys Asp Asp 220 225 230	725
GAC AAG TAC CGA CGT CGG CCG GCC TTG GGT TGG CTG GCC CGG CTG CTA Asp Lys Tyr Arg Arg Arg Pro Ala Leu Gly Trp Leu Ala Arg Leu Leu 235 240 245	773
AGG AGC CGG GCT GGG TCT CGG AAG CGA CCG CTG ACC CTG CTC CAG CGG Arg Ser Arg Ala Gly Ser Arg Lys Arg Pro Leu Thr Leu Leu Gln Arg 250 255 260	821
GCG GGG CTG CTG CTA CTC TTG GGA CTG CTG GGC TTC CTG GCC CTC CTT Ala Gly Leu Leu Leu Leu Gly Leu Leu Gly Phe Leu Ala Leu Leu 265 270 275 280	869
GCC CTC ATG TCT CGC CTA GGC CGG GCC GCA GCT GAC AGC GAT CCC AAC Ala Leu Met Ser Arg Leu Gly Arg Ala Ala Ala Asp Ser Asp Pro Asn 285 290 295	917
CTG GAC CCA CTC ATG AAC CCT CAC ATC CGC GTG GGC CCC TCC TGA Leu Asp Pro Leu Met Asn Pro His Ile Arg Val Gly Pro Ser * 300 305 310	962
GCCCCCTTGC TTGTGGCTAG GCCAGCCTAG GATGTGGGTT CTGTGGAGGA GAGGCGGGGT AATGGGGAGG CTGAGGGCAC CTCTTCACTG CCCCTCTCCC TCAAGCCTAA GACACTAAGA CCCCAGACCC AAAGCCAAGT CCACCAGAGT GGCTCGCAGG CCAGGCCTGG AGTCCCCGTG GGTCAAGCAT TTGTCTTGAC TTGCTTCTC CCGGGTCTCC AGCCTCCGAC CCCTCGCCCC ATGAAGGAGC TGGCAGGTGG AAATAAACAA CAACTTTATT	1022 1082 1142 1202 1242

Figure 2

gb|AA155210|AA155210 mr98e01.r1 Stratagene mouse embryonic carcinoma
(#937317) Mus musculus cDNA clone 605496 5'
Query: 1 MGLCKCPKRKVTNLFCFEH RVNVCEHCLVANHAKCIVQSYLQWLQDSDYNPNCRLCNPL 60
MGLCKCPKRKVTNLFCFEH RVNVCEHCLVANHAKCIVQSYLQWLQDSDYNPNCRLCN PL
Sbjct: 98 MGLCKCPKRKVTNLFCFEH RVNVCEHCLVANHAKCIVQSYLQWLQDSDYNPNCRLCNPL 277

Figure 3

dbj|D75913|CELK111G3F C.elegans cDNA clone yk111g3 : 5' end, single read.
Query: 7 PKRKVTNLFCFEH RVNVCEHCLVANHAKCIVQSYLQWLQDSDYNPNCRLCNIPLASRETT 66
PKRKVTNLFCFEH RVNVCEHCLVANHAKCIVQSYLQWLQDSDYNPNCRLCNIPLASRETT
Sbjct: 1 PKRKVTNLFXYEH RVNVCEELXLVDNHPNCVVQSYLTWLTDQDYDPNCSLCKTTLXEGDTI 180
Query: 67 RLVCYDLFWA CLNERAAQLPRNTAPAGYQCP 98 98 PSCNGPI FPPNQ 109
RL C L HW C +E P TAP GY+CP P C+ +FPP+Q
Sbjct: 181 RLNCLHLLHWKCFDEWXGNFPDTTAPXGYRCP 276 275 PCCSQEVFPFDQ 310

Figure 4

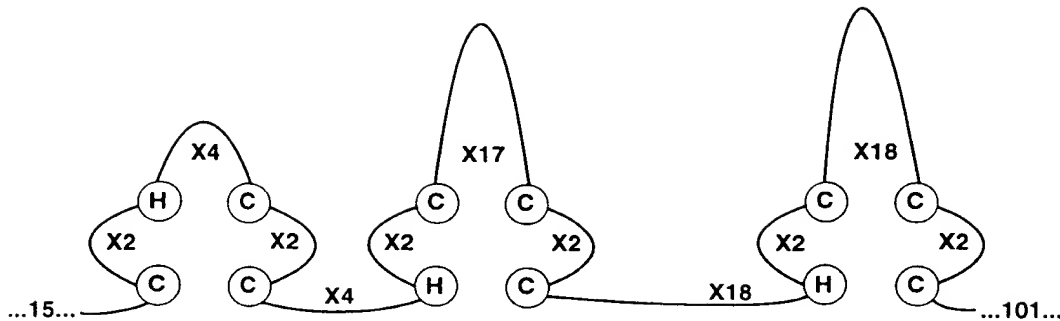


Figure 5

sp|P46580|YLB5_CAEEL HYPOTHETICAL 146.8 KD PROTEIN C34E10.5 IN
CHROMOSOME III gi|500728 (U10402) C34E10.5 gene product
[Caenorhabditis elegans]

Query: 56 CNIPLASREITTRLVCYDLFWAQLNERAQQLPRNTAPAGYQCPSC 100
C+I L ++ + L C LF W C+ E A + + + +CP C
Sbjct: 1222 CSICLENKNPSALFCGHLFCWTCTIQEHAVAATSSASTSSARCPQC 1266

Figure 6

gi|703468 (L29051) homologous to GATA-binding transcription factor
[Schizosaccharomyces pombe]

Query: 35 CIVQSYLQWLQDSDYNPNCRCLCNI 58
C + +W +D NP C C +
Sbjct: 175 CATTNTPKWRRDESGNPICNACGL 198

Query: 162 SSTPGPEEVDSASAAPAFYSQAPRPPASPGRPEQHTVIHMGNPEPLTHAPRKVYDTRDD 221
+S PEE S S S P+ SP + +Q +I P +V + D
Sbjct: 441 ASLLNPEEPSNSDKQPSMSNGPKSEVSPSQSQQAPLIQSSTSPVSLQFPPEVQGSNVDK 500

Query: 222 RTPGLH 227
R L+
Sbjct: 501 RNYALN 506